

CLINICAL RESEARCH PROTOCOL

INITIAL REVIEW APPLICATION

PRINCIPAL INVESTIGATOR (Name, Institute/Branch, Address, Telephone):

PROTOCOL TITLE:

ABBREVIATED TITLE (30 characters or less):

PROPOSED START DATE: _____ END DATE: _____ TOTAL SUBJECTS TO BE ACCRUED (Attach target table for Phase 3-4): _____

COLLABORATING INSTITUTE INFORMATION:

Will subjects participate on the protocol at the NIH CC? ☐ Yes ☐ NoWill subjects participate on the protocol at other sites? ☐ Yes ☐ NoIf yes, are the sites ☐ Domestic ☐ Foreign ☐ Both

Is NIH the coordinating site?

☐ Yes. For each participating site, provide: Institution name, address, investigator(s), indicate if subjects will be recruited and if they are, include a contact name on attached sheet/protocol face sheet.☐ No. Coordinating Site is _____

REQUESTED ACCRUAL EXCLUSION (Check all that apply):

- ☐ None ☐ Asian
☐ Male ☐ Black or African American
☐ Female ☐ White
☐ Children <18 ☐ Hispanic or Latino
☐ American Indian/ Alaskan Native ☐ Native Hawaiian or Pacific Islander

SUBJECT ACCRUAL CHARACTERISTICS:

Minimum Age Permitted _____

Maximum Age Permitted _____

Pediatric ☐ None ☐ <1 Yr. ☐ 1-6 Yrs. ☐ 7-17 Yrs.Protocol involves healthy volunteers? ☐ Yes ☐ NoAre Healthy Volunteers NIH Employees? ☐ Yes ☐ NoSubject Remuneration? ☐ Yes ☐ NoDoes the protocol permit self referral? ☐ Yes ☐ No**NOTE:** The protocol shall include a discussion of the rationale for subject selection/exclusion, including gender and ethnicity of the population at risk, as well as recruitment plans and procedures.

PROTOCOL TYPE: (Check one):

- ☐ Screening
☐ Training
☐ Natural History – Disease Progression
☐ Natural History – Sample/Data Collection or Analysis
☐ Pharmacokinetics/Dynamics
☐ Clinical Trial: Identify Phase (Check one)
☐ Phase 0 ☐ Phase 1 ☐ Phase 1-2
☐ Phase 2 ☐ Phase 3 ☐ Phase 4

If a Phase 3 Clinical Trial, is analysis for sex, racial/ethnic subgroups required according to the NIH Policy and Guidelines on the Inclusion of Women and Minorities as Subjects in Clinical Research? ☐ Yes ☐ No

KEY WORDS (Enter words, not contained in the title, describing the protocol)

1. _____
2. _____
3. _____
4. _____
5. _____

IONIZING RADIATION USE (X-rays, e.g., CT; radioisotopes, e.g. PET; etc.):

☐ None ☐ Medically indicated ☐ Research indicated*

*Complete NIH-88-23a, and attach to this application. Send a copy of entire protocol and NIH-88-23a to Chair, Radiation Safety for concurrent review).

INVESTIGATIONAL NEW DRUG/DEVICE: ☐ None ☐ IND ☐ IDE

FDA No. _____

IND/IDE Name: _____

Sponsor: _____

Who is the manufacturer of the above entity: _____

Does the protocol involve a drug/device/product that may lead to you or the NIH receiving payment and/or royalties?

☐ Yes (Append a statement of disclosure)☐ NoHas the NIH IRP COI Guide been distributed to Non-NIH Investigators? ☐ Yes ☐ No

Based on the criteria at their institution, has a conflict been reported?

☐ Yes. Describe in attached narrative. ☐ No

CONFLICTS OF INTEREST REVIEW:

Date submitted to IC DEC: _____ Date cleared by IC DEC: _____

MEDICAL ADVISORY INVESTIGATOR (if necessary) Initial (Name, Inst/Branch, Telephone, Address, Email and indicate if an NIH employee):

LEAD ASSOCIATE INVESTIGATOR – Initial (Name, Inst/Branch, Telephone, Address, and indicate if an NIH employee):

RESEARCH CONTACT: Initial (Name, Inst/Branch, Telephone, Address, Email and indicate if an NIH employee):

ASSOCIATE INVESTIGATOR(S): Initial (Name, Institute/Branch, Telephone, Address and indicate if an NIH employee)

1. _____
2. _____
3. _____
4. _____
5. _____
6. _____

(Principal Investigator: Be sure to include PRECIS <=400 words as first section of protocol)

SIGNATURE

Principal Investigator

Print/Type Name

Date _____

Send to Accountable Investigator

RECOMMENDATION

Accountable Investigator

Print/Type Name

Date _____

Send to Branch Chief, or CC
Dept. Head of Accountable Investigator

Br. Chief/CC Dept. Head of Acct. Invest.

Print/Type Name

Date _____

Send to Institute/Center Scientific Review
Committee

APPROVALS

For Institute/Center Scientific Review Comm.

Print/Type Name

Date _____

Send to Clinical Director

Clinical Director

Print/Type Name

Date _____

Send to Chair, Institutional Review Board

Chair, For Institutional Review Board

Print/Type Name

Date _____

Send to Office of Protocol Services,
through IRB Protocol CoordinatorPATIENT SAFETY/
RESOURCE REVIEW

Director, Clinical Center

Print/Type Name

Date _____

Return to Office of Protocol Services,
(10/1S231B)

COMPLETION

Protocol Specialist

Date _____

PROTOCOL NO.

Clinical Research Protocol Initial Review Application
NIH-1195 (5-06)

The following data elements are required by the National Library of Medicine for posting on [clinicaltrials.gov](http://www.clinicaltrials.gov) and meets the registration requirements set forth by the International Committee of Medical Journal Editors (ICMJE) for publishing. <http://www.clinicaltrials.gov/>

CONDITIONS: Select up to 5 primary diseases or conditions being studied, using NLM Medical Subject Heading (MeSH) controlled vocabulary. The conditions are used to index studies. <http://www.nlm.nih.gov/mesh/MBrowser.html>

1. _____
2. _____
3. _____
4. _____
5. _____

STUDY TYPE: Nature of the investigation. Select Interventional or Observational, in addition to the most appropriate term describing the protocol for each of the corresponding categories.

<input type="checkbox"/> Interventional Studies	<input type="checkbox"/> Observational Studies
Purpose: Reason for the protocol <input type="checkbox"/> Treatment <input type="checkbox"/> Prevention <input type="checkbox"/> Diagnosis <input type="checkbox"/> Educate/Train Study Design: participant selection <input type="checkbox"/> Randomized Trial <input type="checkbox"/> Non-randomized Trial Masking: knowledge of intervention <input type="checkbox"/> Open <input type="checkbox"/> Single Blind <input type="checkbox"/> Double Blind Control: nature of the interventional control <input type="checkbox"/> Placebo <input type="checkbox"/> Active <input type="checkbox"/> Uncontrolled <input type="checkbox"/> Historical <input type="checkbox"/> Dose Comparison Assignment: intervention groups <input type="checkbox"/> Single Group <input type="checkbox"/> Parallel <input type="checkbox"/> Cross-over <input type="checkbox"/> Factorial <input type="checkbox"/> Expanded Access Endpoint: primary outcome that the protocol is designed to evaluate <input type="checkbox"/> Safety <input type="checkbox"/> Efficacy <input type="checkbox"/> Safety/Efficacy <input type="checkbox"/> Bio-equivalence <input type="checkbox"/> Bio-availability <input type="checkbox"/> Pharmacokinetics <input type="checkbox"/> Pharmacodynamics <input type="checkbox"/> Pharmacokinetics/pharmacodynamics	Purpose: reason for the protocol <input type="checkbox"/> Natural History <input type="checkbox"/> Screening <input type="checkbox"/> Psychosocial Duration of Sampling: protocol sample in <input type="checkbox"/> Longitudinal <input type="checkbox"/> Cross-sectional Selection Method: sample selection <input type="checkbox"/> Targeted Population <input type="checkbox"/> Random Sample <input type="checkbox"/> Case Control Timing: data collection period <input type="checkbox"/> Retrospective <input type="checkbox"/> Prospective <input type="checkbox"/> Both

COMPLETE FOR INTERVENTIONAL STUDIES ONLY

INTERVENTIONS: Provide up to 10 primary interventions identifying a category for each. Category selections are: Drug, Gene Transfer, Vaccine, Behavior, Device, and Procedure.

Category	Intervention	Category	Intervention
Ex. Drug	AZT	Ex. Behavior	Hypnosis
1. _____	_____	6. _____	_____
2. _____	_____	7. _____	_____
3. _____	_____	8. _____	_____
4. _____	_____	9. _____	_____
5. _____	_____	10. _____	_____

OUTCOME MEASURE(S)/ENDPOINT(S): Examples – changes in cardiac output, changes in cognitive function, changes in drug or antibody.

Primary: main outcome representing a primary study question(s). _____

Secondary: outcome(s) of interest to a study, but not representing the primary study question(s). _____